

PCT

NOTIFICATION OF ELECTION

(PCT Rule 61.2)

BOLTON, Robert, John et al

From the	INTERN	ATIONAL	BUREAU
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To:

Commissioner
US Department of Commerce
United States Patent and Trademark
Office, PCT
2011 South Clark Place Room 524
Arlington, VA 22202
ETATS-UNIS D'AMERIQUE
ETATS-LINIS D'AMERIQUE

Date of mailing (day/month/year) 26 October 2000 (26.10.00)	in its capacity as elected Office		
International application No. PCT/AU00/00197	Applicant's or agent's file reference 611547		
International filing date (day/month/year) 17 March 2000 (17.03.00)	Priority date (day/month/year) 17 March 1999 (17.03.99)		
Applicant			

1.	_	ereby notified of its election made: with the International Preliminary Examining Authority on: 12 October 2000 (12.10.00)	
	in a notice effecting	later election filed with the International Bureau on:	
2.	The election X was		
	made before the expiration Rule 32.2(b).	n of 19 months from the priority date or, where Rule 32 appli	es, within the time limit under

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland Authorized officer

Charlotte ENGER

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INTERNATIONAL SEARCH REPORT

International application No. PCT/AU00/00197

A. CLASSIFICATION OF SUBJECT MATTER

Int. Cl. 7: C08G 12/40, 18/83, 59/17; C08F 2/44, 8/44, 283/00, C09D 4/06, 133/02, 133/14, 151/08, 161/32, 163/10

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) C08G 12/40, 59/17, 18/83; C08F 283/00, 220/36, 120/36, 8/44; C09D 133/02, 151/08

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) DERWENT (WPAT and JAPIO)

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
х	US 5847022 A (Y. Yamashina et al.) 8 December 1998 (see entire document)	1-4, 6-9, 15, 19-2
Y	EP 214089 A (Battelle Memorial Institute) 11 March 1987 (see entire document, in particular page 2 line 30 to page 3 line 45)	1, 4, 6, 13, 20-21
Y	US 4297185 A (P. Chevreux et al.) 27 October 1981 (see entire document, in particular column 7 lines 37-43 and column 8 lines 56-68)	1, 4, 6, 13, 20-21

	1 4.4		
*	Special categories of cited documents:	"T"	later document published after the international filing date or
"A"	document defining the general state of the art which is not considered to be of particular relevance		priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"E"	earlier application or patent but published on or after the international filing date	"X"	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an
"L"	document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"Y"	inventive step when the document is taken alone document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is
"O"	document referring to an oral disclosure, use, exhibition or other means		combined with one or more other such documents, such combination being obvious to a person skilled in the art
"P"	document published prior to the international filing date but later than the priority date claimed	"&"	document member of the same patent family

Date of the actual completion of the international search

30 May 2000

Name and mailing address of the ISA/AU

AUSTRALIAN PATENT OFFICE
PO BOX 200, WODEN ACT 2606, AUSTRALIA
E-mail address: pct@ipaustralia.gov.au
Facsimile No. (02) 6285 3929

Date of mailing of the international search report

13 JUNE 2000

Authorized officer

NORMAN BLOM
Telephone No: (02) 6283 2238

INTERNATIONAL SEARCH REPORT

International application No.

PCT/AU00/00197

C (Continuat	tion). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	WO 97/14750 A (W. R. Grace and Co. CONN.) 24 April 1997 (see entire document, in particular pages 9-11 and 14, examples 1, 3, 4, 6, comparative example 2 etc)	1-2, 6-9, 15, 17, 19-21
Y	Polymer Update: Science and Engineering, Edited by W. D. Cook and G. B. Guise, Australian Polymer Science Series Volume 2, Published by the Polymer Division, Royal Australian Chemical Institute, Adams Printers, Victoria Australia, (1989), pages 22-25. (see in particular page 25 last line)	1-2, 6-9, 15, 17, 19-21
X	WO 98/31719 A (EG Technology Partners, LP) 23 July 1998 (see entire document, in particular page 1 lines 20-21, page 3 lines 12-27, page 4, page 8 lines 10-18)	1-4, 6-7, 11, 13-17, 19-21
Α	AU 88546/82 A (Buckman Laboratories, Inc.) 2 June 1983 (see entire document, in particular page 1a lines 10-17 and page 6 lines 17-19, page 3 lines 6-15, page 5 line 26)	1-21
A	US 4451568 A (M. Schneider et al.) 29 May 1984 (see the examples)	1-21
Α	US 4139499 A (K. O. Wade and J. H. Brown) 13 February 1979 (see entire document)	1-21
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INTERNATIONAL SEARCH REPORT Information on patent family members

International application No. PCT/AU00/00197

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent Doo	cument Cited in Search Report			Patent	Family Member		
wo	9714750	AU	73324/96	JP	9124717		
EP	214089	AU	60416/86	CA	1266746	JP	62022864
		US	4766160				
US	5847022	CN	1145926	EP	735426	JP	8325328
US	4297185	СН	652413	EP	10355	US	4355077
US	4451568	wo	8101412	AU	64823/80	DK	3137/81
		EP	29411				
US	4139499	AU	15113/76	CA	1090045	DE	2629146
		FR	2316265	GB	1501307	IL	49920
		IN	1451100	IT	1220912	JP	51017594
		NL	7607168	ZA	7603763		
AU	88546/82	BE	894713	BR	8206542	DE	3237018
		FR	2517314	GB	2112005	IT	8249517
		IT	1157242	JP	58093710	NL	8203877
		NO	823519	NZ	201950	SE	8205445
		ZA	8206951				
wo	9831719	AU	55444/98	EP	952999	GB	9700905

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PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference IRN 611547	FOR FURTHER ACTION			
International Application No.	-	International Filing Date (day/month/year) Priority Date (day/month/year)		
PCT/AU00/00197	17 March 2000		17 March 1999	
International Patent Classification (IPC)	or national classification	on and IPC	1	
	7; C08F 2/44, 8/44, 2	83/00; C09D 4/06, 13	3/02, 133/14, 151/08, 161/32, 163/10.	
Applicant	•			
JARRETT, Kevin George et a	al.			
This international preliminary and is transmitted to the appli	y examination report has icant according to Artic	s been prepared by this Indic 36.	nternational Preliminary Examining Authority	
2. This REPORT consists of a t	otal of 3 sheets, inclu	iding this cover sheet.		
X This report is also acco	mpanied by ANNEXES	, i.e., sheets of the descr	iption, claims and/or drawings which have	
		and/or sheets containing ive Instructions under the	rectifications made before this Authority (see e PCT).	
These annexes consist of a to	tal of 4 sheet(s).			
3. This report contains indications relating to the following items:				
I X Basis of the report				
II Priority				
III Non-establishm	ent of opinion with rega	rd to novelty, inventive	step and industrial applicability	
IV Lack of unity of	invention			
	Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement			
VI Certain docume	nts cited	ts cited		
VII Certain defects	n the international application			
VIII Certain observations on the international application				
Date of submission of the demand		Data of a lation of a	Lagrand	
12 October 2000		Date of completion of t 4 May 2001	ne report	
Name and mailing address of the IPEA/AL	j	Authorized Officer	1000	
AUSTRALIAN PATENT OFFICE			start ()	
PO BOX 200, WODEN ACT 2606, AUS E-mail address: pct@ipaustralia.gov.au	TRALIA		MACON	
Facsimile No. (02) 6285 3929		NORMAN BLOM	(IDON)	
		Telephone No. (02)-62	283 2238	

•	INTERNATIONAL PRELIMINARY EXAMINATION REPORT	PCT/AU00/00197
	3	LETAGOVIOVA
1.	Basis of the rep rt	
1.	With regard to the elements of the international application:	
	the international application as originally filed.	
	X the description, pages 1-19, as originally filed,	
	pages, siled with the demand,	
	pages, received on with the letter of	}
	X the claims, pages, as originally filed,	under Anti-le 10
	pages, as amended (together with any statement) pages, filed with the demand,	under Article 19,
	pages 20-23, received on 16 May 2001 with the	e letter of 16May 2001
	the drawings, pages, as originally filed,	
	pages, filed with the demand,	
	pages, received on with the letter of	
	the sequence listing part of the description:	•
	pages , as originally filed	
	pages, filed with the demand	
	pages , received on with the letter of	
2.	With regard to the language, all the elements marked above were available or fi which the international application was filed, unless otherwise indicated under to These elements were available or furnished to this Authority in the following language.	his item.
	the language of a translation furnished for the purposes of international se	earch (under Rule 23.1(b)).
	the language of publication of the international application (under Rule 4	8.3(b)).
	the language of the translation furnished for the purposes of international and/or 55.3).	preliminary examination (under Rules 55.2
3.	With regard to any nucleotide and/or amino acid sequence disclosed in the in- sequence listing:	ternational application, was on the basis of the
	contained in the international application in written form.	
	filed together with the international application in computer readable for	n.
	furnished subsequently to this Authority in written form.	•
	furnished subsequently to this Authority in computer readable form.	
	The statement that the subsequently furnished written sequence listing do international application as filed has been furnished.	es not go beyond the disclosure in the
	The statement that the information recorded in computer readable form is been furnished	s identical to the written sequence listing has
4.	The amendments have resulted in the cancellation of:	
	the description, pages	·
	the claims, Nos.	
	the drawings, sheets/fig.	
5.	This report has been established as if (some of) the amendments had not go beyond the disclosure as filed, as indicated in the Supplemental Box (
•	Replacement sheets which have been furnished to the receiving Office in response to an report as "originally filed" and are not annexed to this report since they do not contain	
**	Any replacement sheet containing such amendments must be referred to under item ar	

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.
PCT/AU00/00197

v.	Reasoned statement under Ar and explanations supporting s	ticle 35(2) with regard to novelty, invent uch statement	ive step or industrial applicability; citations
1.	Statement		·
	Novelty (N)	Claims 1-22	YES
		Claims none	NO
	Inventive step (IS)	Claims 1-22	YES
		Claims none	NO
	Industrial applicability (IA)	Claims 1-22	YES
		Claims none	NO

2. Citations and explanations (Rule 70.7)

Novelty (N) and Inventive Step (IS): Claims 1-22

The two most relevant documents cited in the International Search Report are WO 98/31719 and US5847022.

WO 98/31719 is distinguished from claim 1-22 for the following reasons.

Although is indicated that lower molecular weight polyamines may be further polymerised to form higher molecular weight polyamines by methods well known in the art, such as by reaction with diepoxides (with ethylene glycol diglycidyl ether being specifically mentioned as suitable diepoxide) (see page 3 lines 12-22), it is to be noted that there is no exemplification of such diepoxide/lower molecular weight polyamine adduct and further, ethylene glycol diglycidyl ether falls outside the scope of claim 1.

US 5847022 is distinguished from claims 1-22 of the present invention because the radiation curable resin is prepared by reacting a tertiary amino group containing polymer (A) with an unsaturated acid (B) to form a protic quaternary ammonium salt, this salt is then reacted with an epoxy compound (C) to form an aprotic hydrophilic quaternary ammonium salt containing resin (see column 7 lines 6-13 etc.). This method of preparation is distinct from that defined by claims 17-20 of the present invention, which require reaction of an unsaturated carboxylic acid with the reaction product of a secondary amine and an epoxy (which process will produce a protic quaternary ammonium salt). Further, claim 1 of the present invention is distinguished from the above citation because it also requires the water soluble amine salt prepolymer to be formed between an unsaturated carboxylic acid and an epoxy-amine adduct formed between an amine and an epoxy of the formula I.

PATENT COOPERATION TREATY PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference IRN 611547	FOR FURTHER See Notification of Transmittal of International Preliminary ACTION Examination Report (Form PCT/IPEA/416).			
International Application No.	International Filing Da	te (day/month/year)	Priority Date (day/month/year)	
PCT/AU00/00197	17 March 2000		17 March 1999	
International Patent Classification (IPC)	or national classification	n and IPC		
Int. Cl. 7 C08G 12/40, 18/83, 59/1	7; C08F 2/44, 8/44, 28	33/00; C09D 4/06, 13	3/02, 133/14, 151/08, 161/32, 163/10.	
Applicant	,			
JARRETT, Kevin George et a	u.		}	
This international preliminary and is transmitted to the applie			nternational Preliminary Examining Authority	
2. This REPORT consists of a to	otal of 3 sheets, includ	ling this cover sheet.		
This report is also accompanied by ANNEXES, i.e., sheets of the description, claims and/or drawings which have been amended and are the basis for this report and/or sheets containing rectifications made before this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT).				
These annexes consist of a tot	al of 4 sheet(s).			
3. This report contains indications relating to the following items:				
I X Basis of the repo	rt			
II Priority				
III Non-establishme	nt of opinion with regard	I to novelty, inventive s	tep and industrial applicability	
IV Lack of unity of	invention			
	V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement			
VI Certain documen	ents cited			
VII Certain defects in	the international applic	ation		
VIII Certain observations on the international application				
Date of submission of the demand		Data of samplation of th		
12 October 2000		Date of completion of th May 2001	e report	
Name and mailing address of the IPEA/AU		Authorized Officer	100	
AUSTRALIAN PATENT OFFICE	1			
PO BOX 200, WODEN ACT 2606, AUST E-mail address: pct@ipaustralia.gov.au	ł		MACION	
Facsimile No. (02) 6285 3929	! T	NORMAN BLOM	TIAC!	
<u> </u>] 7	Telephone No. (02) 62	83 2238	

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International	appli	cation	No
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PCT/AU00/00197

I.	B sis of the report
1.	With regard to the elements of the international application:*
	the international application as originally filed.
	X the description, pages 1-19, as originally filed,
	pages, filed with the demand,
	pages, received on with the letter of
	X the claims, pages, as originally filed,
	pages , as amended (together with any statement) under Article 19,
	pages, filed with the demand,
	pages 20-23, received on 16 May 2001 with the letter of 16May 2001 the drawings, pages, as originally filed,
	pages, filed with the demand, pages, received on with the letter of
	the sequence listing part of the description:
	pages , as originally filed
	pages, as originarly fried
	pages, received on with the letter of
2.	With regard to the language, all the elements marked above were available or furnished to this Authority in the language in which the international application was filed, unless otherwise indicated under this item. These elements were available or furnished to this Authority in the following language which is:
1	the language of a translation furnished for the purposes of international search (under Rule 23.1(b)).
	the language of publication of the international application (under Rule 48.3(b)).
}	the language of the translation furnished for the purposes of international preliminary examination (under Rules 55.2 and/or 55.3).
3.	With regard to any nucleotide and/or amino acid sequence disclosed in the international application, was on the basis of the sequence listing:
	contained in the international application in written form.
	filed together with the international application in computer readable form.
1	furnished subsequently to this Authority in written form.
]	furnished subsequently to this Authority in computer readable form.
	The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished.
	The statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished
4.	The amendments have resulted in the cancellation of:
	the description, pages
	the claims, Nos.
	the drawings, sheets/fig.
5.	This report has been established as if (some of) the amendments had not been made, since they have been considered to go beyond the disclosure as filed, as indicated in the Supplemental Box (Rule 70.2(c)).**
*	Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17).
••	Any replacement sheet containing such amendments must be referred to under item I and annexed to this report

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.
PCT/AU00/00197

	` <u></u>						
v.	Reasoned statement under Ar and explanations supporting s			entive step or industrial applicability; citations			
1.	Statement						
}	Novelty (N)	Claims	1-22	YES			
		Claims	none	NO			
	Inventive step (IS)	Claims	1-22	YES			
		Claims	none	NO .			
	Industrial applicability (IA)	Claims	1-22	YES			
		Claims	none	NO			

2. Citations and explanations (Rule 70.7)

Novelty (N) and Inventive Step (IS): Claims 1-22

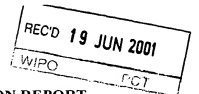
The two most relevant documents cited in the International Search Report are WO 98/31719 and US5847022.

WO 98/31719 is distinguished from claim 1-22 for the following reasons.

Although is indicated that lower molecular weight polyamines may be further polymerised to form higher molecular weight polyamines by methods well known in the art, such as by reaction with diepoxides (with ethylene glycol diglycidyl ether being specifically mentioned as suitable diepoxide) (see page 3 lines 12-22), it is to be noted that there is no exemplification of such diepoxide/lower molecular weight polyamine adduct and further, ethylene glycol diglycidyl ether falls outside the scope of claim 1.

US 5847022 is distinguished from claims 1-22 of the present invention because the radiation curable resin is prepared by reacting a tertiary amino group containing polymer (A) with an unsaturated acid (B) to form a protic quaternary ammonium salt, this salt is then reacted with an epoxy compound (C) to form an aprotic hydrophilic quaternary ammonium salt containing resin (see column 7 lines 6-13 etc.). This method of preparation is distinct from that defined by claims 17-20 of the present invention, which require reaction of an unsaturated carboxylic acid with the reaction product of a secondary amine and an epoxy (which process will produce a protic quaternary ammonium salt). Further, claim 1 of the present invention is distinguished from the above citation because it also requires the water soluble amine salt prepolymer to be formed between an unsaturated carboxylic acid and an epoxy-amine adduct formed between an amine and an epoxy of the formula I.

PATENT COOPERATION TREATY PCT



INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference IRN 611547	FOR FURTHER ACTION		ransmittal of International Preliminary (Form PCT/IPEA/416).
International Application No. PCT/AU00/00197	International Filing Dat 17 March 2000	e (day/month/year)	Priority Date (day/month/year) 17 March 1999
International Patent Classification (IPC)	or national classification	and IPC	
	; C08F 2/44, 8/44, 28	3/00; C09D 4/06, 13	3/02, 133/14, 151/08, 161/32, 163/10.
Applicant	7		
JARRETT, Kevin George et a	1.		
		····	
This international preliminary and is transmitted to the applic			iternational Preliminary Examining Authority
2. This REPORT consists of a tot	al of 3 sheets, includi	ng this cover sheet.	
	e basis for this report and	d/or sheets containing i	ption, claims and/or drawings which have ectifications made before this Authority (see PCT).
These annexes consist of a tota	l of 4 sheet(s).		
3. This report contains indications relating	ng to the following items	:	
I X Basis of the repor	t		
II Priority			$_{\infty}$
III Non-establishmen	t of opinion with regard	to novelty, inventive st	ep and industrial applicability
IV Lack of unity of in	nvention		
	nt under Article 35(2) wanations supporting such		nventive step or industrial applicability;
VI Certain document	s cited		
VII Certain defects in	the international applica	tion	
VIII Certain observation	ons on the international a	pplication	
Date of submission of the demand	D	ate of completion of th	e report
12 October 2000		May 2001	cicport
Name and mailing address of the IPEA/AU		uthorized Officer	$\mathcal{M}_{\mathcal{O}}$
AUSTRALIAN PATENT OFFICE PO BOX 200, WODEN ACT 2606, AUSTI E-mail address: pct@ipaustralia.gov.au Facsimile No. (02) 6285 3929	N	ORMAN BLOM	13 2238



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Internationa	l application	No.

PCT/AU00/00197

I.	Basis of the report
1.	With regard to the elements of the international application:*
	the international application as originally filed.
	X the description, pages 1-19, as originally filed,
	pages , filed with the demand,
	pages, received on with the letter of
	X the claims, pages, as originally filed,
	pages , as amended (together with any statement) under Article 19,
	pages, filed with the demand,
	pages 20-23, received on 16 May 2001 with the letter of 16May 2001
	the drawings, pages, as originally filed,
	pages, filed with the demand,
	pages, received on with the letter of the sequence listing part of the description:
	pages , as originally filed pages , filed with the demand
	pages, filed with the demand pages, received on with the letter of
2.	With regard to the language, all the elements marked above were available or furnished to this Authority in the language in
۷.	which the international application was filed, unless otherwise indicated under this item.
	These elements were available or furnished to this Authority in the following language which is:
	the language of a translation furnished for the purposes of international search (under Rule 23.1(b)).
	the language of publication of the international application (under Rule 48.3(b)).
	the language of the translation furnished for the purposes of international preliminary examination (under Rules 55.2 and/or 55.3).
3.	With regard to any nucleotide and/or amino acid sequence disclosed in the international application, was on the basis of the sequence listing:
	contained in the international application in written form.
	filed together with the international application in computer readable form.
	furnished subsequently to this Authority in written form.
	furnished subsequently to this Authority in computer readable form.
	The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished.
	The statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished
4.	The amendments have resulted in the cancellation of:
	the description, pages
	the claims, Nos.
	the drawings, sheets/fig.
5.	This report has been established as if (some of) the amendments had not been made, since they have been considered to
	go beyond the disclosure as filed, as indicated in the Supplemental Box (Rule 70.2(c)).**
*	Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17).
**	Any replacement sheet containing such amendments must be referred to under item 1 and annexed to this report



International	application No.

PCT/AU00/00197

V.	Reasoned statement under Ar and explanations supporting s	ticle 35(2) with regard to novelty, inventive uch statement	e step or industrial applicability; citations
1.	Statement		
	Novelty (N)	Claims 1-22	YES
		Claims none	NO
	Inventive step (IS)	Claims 1-22	YES
		Claims none	NO
	Industrial applicability (IA)	Claims 1-22	YES
		Claims none	NO

2. Citations and explanations (Rule 70.7)

Novelty (N) and Inventive Step (IS): Claims 1-22

The two most relevant documents cited in the International Search Report are WO 98/31719 and US5847022.

WO 98/31719 is distinguished from claim 1-22 for the following reasons.

Although is indicated that lower molecular weight polyamines may be further polymerised to form higher molecular weight polyamines by methods well known in the art, such as by reaction with diepoxides (with ethylene glycol diglycidyl ether being specifically mentioned as suitable diepoxide) (see page 3 lines 12-22), it is to be noted that there is no exemplification of such diepoxide/lower molecular weight polyamine adduct and further, ethylene glycol diglycidyl ether falls outside the scope of claim 1.

US 5847022 is distinguished from claims 1-22 of the present invention because the radiation curable resin is prepared by reacting a tertiary amino group containing polymer (A) with an unsaturated acid (B) to form a protic quaternary ammonium salt, this salt is then reacted with an epoxy compound (C) to form an aprotic hydrophilic quaternary ammonium salt containing resin (see column 7 lines 6-13 etc.). This method of preparation is distinct from that defined by claims 17-20 of the present invention, which require reaction of an unsaturated carboxylic acid with the reaction product of a secondary amine and an epoxy (which process will produce a protic quaternary ammonium salt). Further, claim 1 of the present invention is distinguished from the above citation because it also requires the water soluble amine salt prepolymer to be formed between an unsaturated carboxylic acid and an epoxy-amine adduct formed between an amine and an epoxy of the formula I.

Claims:

1. An ultraviolet light curable resin composition comprising a water soluble amine salt prepolymer formed between an unsaturated carboxylic acid and an oligomer having at least one amine group selected from the group consisting of urea formaldehyde resins, melamine formaldehyde resins, amine polyisocyanate adducts, Michael adducts of a secondary amine and acrylate and/or methacrylate compounds and epoxy-amine adducts formed between an amine and epoxy of formula:

10

5

$$\begin{array}{c}
O \\
CH_2 \longrightarrow CHCH_2 \longrightarrow O-A-O-CH_2CHCH_2 \\
\hline
O-A-O-CH_2CHCH_2 \\
\hline
n
\end{array}$$

wherein n is from 0 to 10 and A comprises a diradical selected from the group consisting of aromatic, substituted aromatic and the diradical of formula II:

$$R$$
 0 11

wherein R is an alkylene group of 1 to 4 carbon atoms.

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- 2. A resin composition according to claim 1 comprising an aqueous solution containing a weight ratio of amine salt prepolymer to water in the range of from 1:4 to 20:1.
- 25 3. A resin composition according to claim 2 wherein said weight ratio is in the range of from 3:2 to 9:1.
 - 4. A resin composition according to claim 1 which is curable by UV and contains less than 0.5% by weight based on the weight of the resin component of a UV initiator.

- 5. A resin composition according to claim 1 which is curable by UV in the absence of photoinitiator.
- 6. A resin composition according to claim 1 wherein the oligomer having at
 5 least one amine group is an epoxy-amine formed between a secondary amine and an epoxide of formula 1:

$$\begin{array}{c}
O \\
CH_2 \longrightarrow CH CH_2 \longrightarrow O-A-O-CH_2CH CH_2 \longrightarrow O-A-O-CH_2 CH \longrightarrow CH_2
\end{array}$$

wherein n is from 0 to 10 and A is a diradical of formula II:

wherein R is an alkylene group of 1 to 4 carbon atoms.

- 7. A resin composition according to claim 6 wherein the epoxide of formula 15 1 is a product of epichlorohydrin and bisphenol A.
 - 8. A resin composition according to claim 6, wherein the amine adduct is formed using a molar ratio of amine to epoxide compound in the range of from 0.5:1 to 2.05:1.
 - 9. A resin composition according to claim 6 wherein the molar ratio of amine to epoxide compound is 0.95:1 to 1.6:1.
- 10. A resin composition according to claim 1 wherein the oligomer having at least one amine group is prepared from melamine, formaldehyde and/or glyoxal and optionally one or more alcohols selected from the group consisting of C₁ to C₆ alkanols, C₁ to C₆ alkyl ethers of C₁ to C₆ alkylene glycols and C₁ to C₆ alkylene glycols.

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- 11. A resin composition according to claim 1 wherein the oligomer having at least one amine group is a urethane-amine adduct prepared by reaction of a secondary amine with a polyisocyanate.
- 5 12. A resin composition according to claim 1 wherein the urethane amine adduct is formed from a molar ratio of amine to polyisocyanate of from 0.90:1 to 1.6:1.
- 13. A resin composition according to claim 1 wherein the oligomer having at least one amine group is a Michael adduct formed between a secondary amine and an unsaturated compound selected from the group consisting of monomers and prepolymers comprising a plurality of unsaturated groups including at least one acrylate or methacrylate group.
- 15 14. A resin composition according to any one of claims 6 to 9 wherein the secondary amine is amine of formula IV:

$$R \longrightarrow N \longrightarrow R^1$$
 IV

- wherein R and R¹ are independently selected from straight and branched chain aliphatic of up to 6 carbon atoms optionally substituted by hydroxy and mixtures thereof.
- 15. A resin composition according to claim 1 wherein the unsaturated acid is selected from the group consisting of acrylic acid, methacrylic acid, crotonic acid, citraconic acid, sorbic acid, fumaric acid and mixtures of two or more thereof.
- 16. A resin according to claim 1 wherein the unsaturated acid is selected 30 from the group consisting of acrylic acid and methacrylic acid.
 - 17. A method of preparing a radiation curable composition comprising forming an amine adduct by reaction of a secondary amine and an epoxy and

reacting the amine adduct with an unsaturated carboxylic acid to form a water soluble salt.

- 18. A method according to claim 17 wherein the amine adduct and unsaturated carboxylic acid are reacted in the presence of water to provide an aqueous solution of a salt formed between the epoxy amine adduct and unsaturated carboxylic acid.
- 19. A method according to claim 17 wherein the reaction of the secondary10 amine and epoxy is exothermic and the reaction mixture is diluted with water during the exotherm.
 - 20. A method according to any one of claims 17 to 19 wherein water is added to provide a concentration of from 20 to 95%.
 - 21. A method of forming a coating on a substrate comprising applying to the substrate a layer of an aqueous solution of a radiation curable resin according to claim 1 and subjecting the layer of said aqueous solution to ultraviolet radiation to cure the layer.
 - 22. The method according to claim 21 wherein the composition is applied at a thickness of up to 200 microns.

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(54) Title: RADIATION CURABLE RESIN COMPOSITION

(57) Abstract

A radiation curable resin composition comprising a water soluble amine salt prepolymer formed between an oligomer or compound having at least one amine group and an unsaturated carboxylic acid and use of this composition as a surface coating or ink.

PCT/AU00/00197

Radiati n Curable Resin Composition

The present invention relates to radiation curable resin compositions and to method of using such compositions.

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The use of ultraviolet (UV) radiation or electron beam (EB) radiation as the energy source for cross linking functional monomers, oligomers and polymers is the basis of important commercial processes with broad applicability in photo imagery coatings inks and adhesive.

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The most predominant type of resin systems used contain an oligomer such as epoxy acrylate, urethane acrylate or unsaturated polyester, a monomer diluent which is a polyunsaturated acrylic or vinyl monomer and an initiator.

The radiation curing processes, particular those involving UV radiation, generally rely on a polymerisation initiator which in the presence of the appropriate radiation generates activated species which initiate the polymerisation reactions and produce curing. Examples of photoinitiators used in UV curable composites include benzoin ethers such as benzyl methylether,
 hydroxydialkylacetophenones such as a α-hydroxy-α,α-dimethylacetophenone and 1-benzoylcyclohexanol, acylphosphine oxides, cyclic benzoin ethers, benzophenone, Michlers ketone and ketocoumarins.

The currently used radiation curable resin compositions are relatively expensive when compared with other coating systems due to the relatively high cost of resins, monomers and photoinitiators.

In many instances it is also difficult to achieve effective surface cure of the resins. Oxygen tends to inhibit polymerisation at the surface leading to undesirable surface tackiness. The use of certain photoinitiator combinations such as benzophenone and amines have been used to counteract this problem but may lead to undesirable odour or colouration. Inert gases have been used to reduce oxygen-inhibition but they are a further expense and are impractical in many commercial environments.

PCT/AU00/00197

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The existing compositions are also completely polymerisable to avoid the use of solvents or diluents. This makes it difficult in many instances to formulate semigloss and flat coatings or films of low film weight.

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Furthermore many of the low viscosity, low molecular weight monomers have a high draize value and their use is minimized for health reasons.

We have now developed versatile radiation curable resin compositions which significantly reduces and in many cases eliminates the need for addition of a photoinitiator and which may address one or more further shortcomings of the prior art described above.

Accordingly we provide a radiation curable resin composition comprising a water soluble amine salt prepolymer formed between an oligomer having at least one amine group and an unsaturated carboxylic acid.

One of the significant advantages of the prepolymers used in the composition of the invention is their water solubility. The prepolymer may generally be diluted in water to at least 50% by weight permitting their use not only in high gloss but also matt or semi gloss films.

The weight ratio of the amine salt prepolymer to water in the resin composition of the invention is typically in the range of from 1:4 to 20:1 more preferably from 3:2 to 9:1 and most preferably from 3:2 to 8:1.

Despite their solubility in water the resins will generally polymerise on exposure to radiation after application to a substrate forming tough water resistant coatings without the need to take any special steps to remove water although dying may be promoted using known methods if desired.

The oligomer component of the salt may be selected from a range of oligomers. Typical example of suitable classes of oligomers include epoxy-amine adducts, amino resins such as urea formaldehyde and melamine formaldehyde type

PCT/AU00/00197

3

resins, amine-polyisocyanate adducts, and Michael adducts of an aliphatic amine and polyacrylate or polymethacrylate compound.

The preferred oligomer components are selected from epoxy-amine and amino resins.

The most preferred prepolymers for use in the radiation curable composition of the invention are a salt of an epoxy-amine adduct and the unsaturated acid.

The epoxy resins from which the adduct may be formed are known in the art.

One class of suitable epoxies is of the formula I:

n is from 0 to 10 and

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A is a diradical selected from the group consisting of aliphatic, substituted aliphatic, aromatic, substituted aromatic, cycloaliphatic, substituted cycloaliphatic alkylene dicarbonyl and the diradical group of formula II:

wherein R is an alkylene group of 1 to 4 carbon atoms. Preferred group A include C₂ to C₂₂ alkyl dioxybenzene and C₄ to C₂₂ alkylene dicarbonyl and groups of formula II. Most preferably A is of formula II.

The most preferred epoxy is a condensation product of bisphenol A and epichlorohydrin.

The amine will generally react with the terminal epoxy groups to form one or more terminal amine groups depending on the stoichiometry of the reaction. In cases where the stoichiometry used does not produce a reaction with all epoxy groups the remaining groups may be reacted to provide other functional groups

PCT/AU00/00197

such as ester groups by reaction with unsaturated acids or ethers by reaction with alcohols or alcohol groups by reaction with water.

The stoichiometry of the epoxy amine adduct is preferably in the range of from 0.5 to 2.05 mole of amine compound per mole of epoxy compound, more preferably from 0.95 to 1.6 and most preferably from 1.0 to 1.5.

The reaction between the epoxy and amine is exothermic and if allowed to proceed may produce an undesirably viscous resin. It is therefore preferable to control the reaction temperature and also quench the reaction by combining the reaction product with water and preferably stirring the aqueous mixture. The unsaturated acid is preferably added to the aqueous mixture and generally produces a clear resin solution. In this case excess unreacted epoxy groups may react with water to form diols.

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A further embodiment of the invention uses amino resins. The preferred amino resins are based on the reaction products of formaldehyde and urea or melamine. Formaldehyde reacts with urea to form resins which are generally bifunctional. Melamine on the other hand reacts with formaldehyde to form trifunctional resins.

Urea and melamine with formaldehyde and glyoxal are the basic materials but melamine may be partly or completely replaced by benzoguanamine and acetoguanamine. Urea/formaldehyde and/or melamine/formaldehyde are the preferred amino resins. Further examples of preferred type of amino resins include ethylene urea, a resin based on dimethylolethylene urea is prepared from urea, ethylene diamine and formaldehyde; propylene urea-formaldehyde, a resin prepared from urea, 1,3-diamino-propane and formaldehyde; triazone resin made from urea, formaldehyde and a primary aliphatic amine such as hydroxyethylamine, uron resins, mixtures of a minor amount of melamine resin and uron (predominantly N,N'-bismethoxymethyl) uron plus about 25% methylated urea formaldehyde; glyoxal resin prepared from urea and/or melamine, glyoxal and formaldehyde and one or more alcohols such as ROH where R is alkyl or an alkyl hydroxy alkyl or alkyl ether of ethylene glycol; 14/09 '01 FRI 11:46 FAX 613 96 2294

WO 00/55228

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PCT/AU00/00197

melamine-formaldehyde which include the dimethyl ether of trimethylolmelamine; and methylol carbamate resins, derivatives made from urea and an alcohol, ROH wherein R is an alkyl, hydroxy alkyl or alkyl ether of ethylenglycol.

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The more preferred amino resins include melamine and formaldehyde and optionally glyoxal and/or urea.

The preferred melamine formaldehyde resins are of the glyoxal type and are prepared from melamine, formaldehyde, glyoxal and preferably further including one or more alcohols selected from the group consisting of C₁ to C₆ alkanols, C₁ to C₆ alkylenes of C₁ to C₆ alkylene glycols and C₁ to C₆ alkylene glycols. To provide good shelf stability it is particularly preferred to include an alcohol. We have also found the efficiency of cure of melamine formaldehyde resins may be improved in many cases by using an acid buffering agent such as phosphoric acid to preferably provide a pH of less than 5.

The oligomer compound of the composition of the invention may be a urethane amine adduct. The amine adduct may be prepared by reaction of an amine with a polyisocyanate. The polyisocyanate may be a simple polyisocyanate such as toluene diisocyanate, diphenylmethane-4,4-diisocyanate or hexamethylene diisocyanate or may be prepared by reaction of a polyol and polyisocyanate. Examples of suitable polyol components include polyester polyols, polyether polyols and polyester polyether block copolymers.

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The molar ratio of amine to polyisocyanate used in forming the adduct is typically from 0.90 to 1.6 and most preferably from 1 to 1.5.

Urea groups may be prepared from isocyanate groups of a urethane prepolymer by reaction with water. The preferred urethane amine adduct is based on an aromatic isocyanate.

The oligomer component of the composition of the invention may be a michael adduct formed by reactions of amine with a multifunctional acrylate or

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PCT/AU00/00197

methacrylate monomer or prepolymer or mixture of two or more thereof. The michael adduct is preferably formed between a secondary amine and an unsaturated carboxylic acid selected from monomers and oligomers containing a plurality of unsaturated groups including at least one acrylate or methacrylate group. Examples of suitable multifunctional acrylates and methacrylates include urethane acrylates, urethane methacrylates, diisocyanates, polyol diacrylates and dimethacrylates, alkylene diacrylates, alkylene dimethacrylates, oligoester diacrylates and oligoester dimethacrylates wherein the oligoester portion may be formed by reaction of adipic, phthalic, isophthalic acid or their anhydrides with polyols such as alkylene glycol or polyalkylene glycol as well as triols such as trimethylol propane and tetrols such as pentaerylthritol.

Examples of michael adducts are described in US Patent 3844916 (Gaske) and will preferably be formed by reaction of a secondary amine and multifunctional acrylate/methacrylate.

The Michael adducts will generally include an amino group of formula III as a result of the addition:

wherein R and R¹ are preferably as defined below for the amine of formula IV and R² is methyl or hydrogen.

The amine group or groups which are present in the prepolymer component of the composition of the invention are preferably selected from secondary amines, secondary alkanolamines and mixtures of two or more compounds from these groups. The preferred amines are of formula IV:

$$R \longrightarrow N \longrightarrow R^1$$
 IV

7

PCT/AU00/00197

wherein R and R^1 are independently selected from the group consisting of straight and branched chain aliphatic of up to 10 carbon atoms optionally substituted by hydroxy and preferably R and R^1 are C_1 to C_6 alkyl optionally substituted by hydroxy. The hydroxy substitution is preferably on a primary carbon atom which is not directly adjacent an amino nitrogen. Particularly preferred R and R^1 are ethyl, 2-hydroxy ethyl, propyl, 3-hydroxypropyl. Most preferably the amine is diethylamine or diethanolamine.

It is preferred that the amine component is substantially free of primary amine and preferably it-contains less than 5% by weight of primary amine. This is particularly preferred where the oligomer is an amine adduct of an epoxy or polyisocyanate. The presence of amine compositions containing at least 95% secondary amine and preferably at least 98% produces lower viscosity resins which tend to have better long term stability.

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The oligomer component of the radiation curable resin composition of the invention is in the form of a salt with an unsaturated carboxylic acid. The preferred unsaturated acids are αβ unsaturated carboxylic acid particularly those selected from the group consisting of acrylic acid, citraconic acid, sorbic acid, fumaric acid and mixtures of two or more thereof. The preferred unsaturated acids are compounds of the formula:

$$CH_2 = C - C - OH$$

$$\downarrow R$$

wherein R is hydrogen C₁ to C₄ alkyl and preferably hydrogen or methyl. The preferred unsaturated acids are thus acrylic acid and methacrylic acid. Mixtures of two or more unsaturated acids may also be used.

The molecular weight of the oligomer resins used in the composition of the invention may vary significantly depending on their proposed use and concentration. Typically the molecular weight of the oligomer salt is in the

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PCT/AU00/00197

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range of from 500 to 20,000 more preferably from 700 to 10,000 and most pref rably from 1000 to 3000.

The resins compositions of the invention may be used in aqueous formulations to form coatings by UV radiation curing. No added initiators are generally required to obtain efficient UV curing although they may be used if desired. Typically the composition will contain less than 0.5% by weight of added initiators and are most preferably free of added photoinitiators. Polymerisation under UV occurs rapidly without generally needing to dry the water used in forming the salts. The UV cured film forms a tack free surface at the air interface.

The composition of the invention is preferably a coating composition or an ink. The composition may be formulated with suitable additives. Examples of suitable additives include surfactants, antifoam agents, inhibitors, pigments, fillers and other resins. Solvents may be included if desired yet are generally not required.

Surfactants may be used where the composition is to be used as a coating for greasy surfaces such as for overcoating inks. Examples of suitable surfactants include silicone derivatives, vinyl silanes and fluorinated wetting agents such as fluoroalkyl ethers.

Inhibitors may be used to inhibit gelling of the composition during storage. Examples of inhibitors include hindered amines such as diphenylamine, phenothiazine and di-p-fluorophenylamine; phenolic and quinone/hydroquinone compounds such as hydroquinone, mono-tertbutylhydroquinone, benzoquinone, 2,5-diphenyl-p-benzoquinone, p-methoxyphenol and 2,6-di-tert-butyl-cresol Inhibitors are generally used in amount of from 0.01 to 1% and preferably 0.01 to 0.5% by weight.

In a further aspect the invention provides a method of preparing a radiation curable composition comprising forming an amine adduct by reacting of a

14/09 '01 FRI 11:47 FAX 613 96 2294

WO 00/55228

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9

PCT/AU00/00197

secondary amine with an epoxy and reacting the amine adduct with an unsaturated carboxylic acid to form a water soluble salt.

The amine adduct and unsaturated carboxylic acid are generally reacted in the presence of water. The reaction which takes place between a secondary amine and epoxy or polyisocyanate is exothermic and we have found that particularly good results are obtained if the reaction mixture of the epoxy or polyisocyanate and secondary amine are diluted with water during the exotherm. This may be done by adding an excess of water or by pouring the reaction mixture into water-

It is surprising to find that the compositions of the invention are generally polymersable by standard UV radiation equipment without the need to use a photoinitiator. The most preferred resins from this point of view are the unsaturated acid salts of epoxy-amine adducts and amine resins. Isocyanate adducts are in some cases less sensitive and may require addition of small amount of for example up to 0.5% by weight of photoinitiator. The reduction or elimination of the photoinitiator component significantly reduces the cost of compositions of the invention when compared with traditional UV curable coatings. Furthermore the component used in preparing the compositions of the invention may be chosen from inexpensive resins, the components of which are readily available. The relatively expensive monomer components required in many prior art compositions can be avoided.

The sensitivity of the compositions of the invention can be compared for different resins and with prior art compositions using the following curing test.

In the curing test a coating film of 5 microns is drawn down on a substrate and is exposed to ultraviolet light of 25 millijoules per cm². The amount of energy may be measured using a suitable radiometer such as an EIT Uvicure Radiometer Model M365. The lamp which we used in our test was a mercury lamp of 80 w/cm at half power (40 w/cm) and we used a belt speed of 35 metres per minute.

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PCT/AU00/00197

The compositions of the invention will generally cure under the above test conditions with addition of less than 0.5% based on resin component of initiator and the compositions of the invention will in most cases require no added photoinitiator. In contrast compositions of the prior art generally require 5% photoinitiator based on the resin component.

Although the resin compositions generally cure without addition of a photoinitiator the compositions may, if desired contain conventional initiators. Typically when used the photoinitiator will be present in an amount of less than 0.5% by weight based on the total weight of the resin component.

The resin composition is typically prepared as an aqueous solution containing at least 10% by weight oligomer salt. The aqueous solution will typically contain in the range of from 20 to 95% by weight of the resin salt preferably form 50 to 85% and most preferably in the range of from 60 to 80% by weight of resin.

A further significant advantage of compositions of the invention is that their cure is generally not significantly inhibited by oxygen when cured in air. As previously indicated the curing of conventional compositions are inhibited by oxygen in air making it difficult to rely on UV curing for preparing coatings without excluding air by using an inert gas or specific initiator combinations which further increase costs.

The ability to prepare the resins at a range of concentrations makes them particularly versatile in controlling gloss levels. Flat films may be prepared using a relatively dilute composition such as from 40 to 65% by weight resin in water. Gloss films on the other hand may contain 65 to 90% by weight resin in water.

Prior art compositions made it difficult to achieve the same properties without the compositions being undesirably viscous or containing a high proposition of relatively expensive monomer diluent.

11

PCT/AU00/00197

Even though the compositions of the invention are prepared and used as an aqueous solution they dry to form a water resistant film which is durable and resistant to solvents.

The compositions of the invention are also relatively non hazardous to skin contact and are essentially odour-free.

Each of these advantages alone has merit and collectively the advantages of compositions of the invention constitute a major commercially attractive advance to the field of radiation cured coatings.

The invention will now be described with reference to the following examples. It is to be understood that the examples are provided by way of illustration of the invention and that they are in no way limiting to the scope of the invention.

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Examples 1 to 7

These examples demonstrates preparation of compositions of the invention in which a portion of the epoxy groups are pre-reacted with acrylic acid prior to addition of a secondary amine to form the epoxy amine adduct and subsequently formation of the amine/acrylic acid salt.

To 0.1 mole (36gm of bisphenol A diglycidyl ether (Araldite 6010) is added varying amounts of acrylic acid together with 0.1g hydroquinone monomethyl ether inhibitor and 0.5g catalyst. The mixtures are reacted at 100°C for 40 minutes and then cooled to 95°C, and diethanolamine is added over several minutes, allowing the subsequent exotherm to proceed to 140-150°C with no external heat. The product is allowed to cool to 90°C, and water is then added with rapid stirring producing a white stable dispersion. Acrylic acid is then added slowly with continuous stirring to solublise the resin solution.

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After 15 minutes, the solution is allowed to cool, inhibitor added and transferred to a dark glass container.

PCT/AU00/00197

12

1	36g (0.1 mol)	5.8g (0.08 mole)	12.6g (0.12 mol)	20g	1 0 g	>1p
2	36g (0.1 mol)	5,6g (0.09 mol)	11.5g (0.11mol)	20g	79	>1p
3	36g (0.1 mol)	7.2g (0.10 mol)	10.5g (0.10 mol)	20g	6g	420cp
4	36g (0.1 mol)	7.9g (0.11 mol)	9.5g (0.09 mol)	20g	6g	980cp
5	36g (0.1 mol)	8.6g (0.12 mol)	8.4g (0.08 mol)	20g	5g	110cp
6	36g (0.1 mol)	9.4g (0.13 mal)	7.4g (0.07 mol)	20g	6g	620cp
7	36g (0.1 mol)	10.1g (0.14 mol)	6.3g (0.06 mol)	209	4g	>1p

Each sample was then coated onto calendared paper at 5 microns using a wire wound applicator, and passed under UV light at 15m/minute. Each film was examined for gloss, surface tack, hardness, water and MEK resistance. The results were as follows:

Examples 3, 4 and 5 all cured after one pass to give satisfactory films.

Examples 1 and 2 were more difficult to apply due their viscosity and required two passes to cure.

Examples 6 and 7 were slightly more turbid and produced dull films.

Example 8

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This example and examples 9 to 21 below demonstrate the preparation of an epoxy amine resin which is reacted with an unsaturated acid to form a resin salt in accordance with the invention. One mole of bisphenol A diepoxide resin was reacted with at least 1 mole of diethanolamine to form an adduct. Allowing the exotherm of this reaction to take its course over 5-10 minutes. The composition was diluted with water or quenched into water to form a stable dispersion of the adduct. To this dispersion, 2 moles of acrylic acid or methacrylic acid were added to produce a cationic resin solution, with very similar properties of those resins formed from epoxy acrylate half ester described in Examples 1 to 7. These resins have low odour and colour and will photopolymerise, when exposed to UV light, to a hard tack-free state.

The resin is of low viscosity and permits solids contents of at least 80%, with as little as 20% water, and have moderately low viscosity. The overall reaction

PCT/AU00/00197

13

time can be accomplished within 1 hour, by avoiding the longer ester formulation of the method described in Examples 1 to 7. The resins of this example appears to show reduced tendency to auto-polymerise.

5 Example 9

Bisphenol di-glycidyl ether resin (1 mole) 360g is mixed with diethanolamine (1 mole) 105g at room temperature, and heated with stirring to 70°C, when it forms a clear homogeneous mixture. At this stage an energetic reaction takes place and the external heat is removed. The resultant exotherm increases the temperature rapidly to 140-150°C with a corresponding reduction in viscosity of the clear fluid. This is rapidly poured into (10 moles) 180g of cold water, with fast stirring to produce a white dispersion, with corresponding increase in temperature of the resin mixture. Acrylic acid (2 moles) 144g, is added over several minutes and a clear resin solution results. The resin solution is allowed to cool, with continuous stirring. The resultant resin is a clear, colourless, low viscosity oligomer solution with almost no odour and a pH of 4-6. When cast as a thin film onto a metal or paper substrate, and exposed to UV light for a few seconds, the resin rapidly produces a hard, tack-free and odour-free clear glossy film, with good water and solvent resistance.

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Examples 10-15

The procedure of Example 8 was followed with the exception that water was added to the resin at the stage of exotherm in which the resultant reflux of water assisted in controlling the heat and provided the dispersion prior to salt formation. Different amines and unsaturated acids were also examined.

	Moles	Moles	Moles	Moles	Moles	Moles
BPDGE (Araldite 6010)	1.0	1.0	1.0	1.0	1.0	1.0
Diethanolamine	1.0	1.0	1.3	1.6	1.0	1.0
Water	10.0	10.0	10.0	10.0	10.0	10.0
Acrylic acid	1.0	2.0	2.0	2.0	-	-
Methacrylic acid		-			1.0	2.0



PCT/AU00/00197

14

Each of these resin solutions was a clear low viscosity solution and produced a hard glossy film after exposure to UV radiation. Example 13 did show a tendency to water sensitivity at the higher ratio of diethanolamine.

5 Examples 16-21

1. 15.7 (v) 1.0 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1		(F				
	Moles	Moles	Moles	Moles	Moles	Moles
BPDGE (Araldite 6010)	1.0	1.0	1.0	1.0	1.0	1.0
Diethanolamine	1.0	1.0	1.3	1.6	1.0	1.0
Diethylamine		1.0	0.7	0.4	-	
Urea		-	-	-	0.5	1.0
Water	10	10	10	10	10	10
Acrylic acid	2.0	2.0	2.0	2.0	2.0	2.0

These resins displayed differences in viscosity, but each was photocurable and produced satisfactory films.

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Examples 22-28

These examples examine preparation of compositions of this invention using melamine formaldehyde based resins.

15 **Example 22**

To 324g of 37% formalin (4 moles) is added 126g of melamine (1 mole) at room temperature and heated with stirring, to 70°C and held until clear, about 40 minutes. The solution is cooled to 45°C and 145g of 40% glyoxal (1 mole) is added with constant stirring, and held for a further 30 minutes. A clear pale amber solution results, to which is added 144g of acrylic acid (2 moles), and allowed to cool. The resultant solution is a very pale, clear and odour-free solution which when cast onto a metal or paper substrate and exposed to UV radiation, will cure rapidly to a hard, tack-free and odour-free clear film, with good water and solvent resistance.

14/09 '01 FRI 11:49 FAX 613 96 2294

WO 00/55228

PCT/AU00/00197

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The procedure of Example 22 was repeated using various proportions of components and alcohols as shown in the table below.

	Moles	Moles	Moles	Moles	Moles	Moles
Formalin 37%	3	3	3	3	3	3
Giyoxan 40%	1	1	1 1	1	1	1
Melamine	1	1	1	1	1	1
Ethanol	-	2	-	-	2	1
n-Propanol		-	2	-	-	2
Hydroxy ethyl acrylate	-	1 -	1 =	1	-	-
Acrylic acid	2	2	2	 - -	1	2

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Example 29

In a similar manner, to 324g of 37% formalin (4 moles) is added 126g of melamine and also 240g of n-propanol (4 moles) and heated with stirring to 65°C and held for 30 minutes. A clear solution results which smells faintly only of propanol, to which is added 144g of acrylic acid (2 moles), and the stirring continued for a further 30 minutes at 50°C, then cooled. This resin solution behaves in the same way as Example 12, will UV cure to a clear glossy film with faint odour of propanol. It does appear to be slightly more flexible in thicker films than Example 22 films.

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Example 30-33

Resins were prepared according to Example 29 but replacing n-propanol with methanol (Example 30), ethanol (Example 31), butanol (Example 32) and isopropanol (Example 33). Each resin responded to UV radiation. The isopropanol resin was the least reactive thought to be due to steric hindrance. The resin based on n-butanol had a stronger residual odour.

Example 34

Other resins were made from monofunctional epoxides, by similar reactions to that described in Example 1 to 7 except that the full diethanolamine adduct was

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PCT/AU00/00197

formed, followed by salt formation with acrylic acid. Six typical examples were made from cresol glycidyl ether, butanediol glycidyl ther, C12-14 alkyl glycidyl ether, neodecanoic acid diglycidyl ester, as well as butanediol di glycidyl ether and an alicyclic diepoxy carboxylate. These all exhibited some of the desired properties as indicated above, yet were all of lower viscosity and less water resistant.

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Example 35

To 65g(0.8 moles)of 37%formalin was added 24.5g urea(0.4 moles)and 48g npropanol (0.8 moles)and heated with stirring to 60 degrees C, and held for 20
minutes. Heat was stopped and acrylic acid 60g (0.8 moles)slowly added over
3 minutes with continuous stirring. At about 40 degrees, phosphoric acid 1 g
was added and then cooled to room temperature. The resulting clear liquid resin
was found to UV cure to a soft film which continued harden on further
irradiation.

Similarly, resins were prepared by replacing propanol with 36g of ethanol which gave a clear resin product with similar properties.

Example 36

To 33g (0.4 moles) 37% formalin was added 16g (0.13 moles) melamine and 2.0 g diethanolamine heated with stirring to 60 degrees C and held for 10 minutes, 16g (0.25 moles) n-propanol was added and held at 60 degrees for a further 20 minutes. The resin was cooled to 30 degrees and acrylic acid 18g (0.25 moles) and phosphoric acid 2g added and stirred until cold. The resultant clear resin cures to a dry glossy film under UV radiation.

A similar batch was prepared except acrylic acid was increased by 50% to 27g (0.38 moles) and no phosphoric acid. This resin was very similar and UV cured as before.

PCT/AU00/00197

A third resin was prepared by replacing propanol with 14 g (0.25 moles) of ethanol and then adding 18 g acrylic acid and 2g phosphoric acid. This resin behaved as in the other cases.

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5 Example 37

Using the same general procedure urea (1 mole, 60g) was reacted with Glyoxal (1 mole, 145g of a 40% solution) and then acrylic acid (2 moles, 144g) added to form a salt. The resultant solution was capable of UV cure but the so formed film was water sensitive.

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Example 38

One mole of propylene oxide (58g) was carefully added to one mole (105g) of diethanolamine, and allowed to cool. Water (20g) was added and then one mole (72g) of acrylic acid. The adduct was tested as before and found to be water soluble, low odour and colour, low viscosity but only partially cures under UV radiation.

Example 39

This example demonstrates preparation of compositions of the invention from michael adducts. One mole of tri propylene glycol diacrylate (300g) was mixed with one mole (105g) of diethanolamine and heated to 105°C and held for 30 minutes. The product was then cooled to 90°C and 90g water added with stirring. 1 mole (72g) of acrylic acid was slowly added. On testing for cure, this resin was found to cure only with 1-2% of photoinitiator (Irgacure 184).

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Example 40

To the resin of Example 3 was added 20% of aqueous pigment dispersions based on carbon black, diarylide yellow and phthalocyanine blue were added with stirring. These pigmented compositions were stable and when cast out as films and exposed to UV radiation, cured in a similar manner to the unpigmented resins previously described. The presence of pigment adversely affects the rate of cure, and small additions of photoinitiator may be used to accelerate the cure.

18

PCT/AU00/00197

Example 41 and Comparativ Example A

The composition of Example 20 was formed into a coating composition for use as a high gloss overprint varnish having the following compositions

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Example 41

	100.0
Water	15.0
(50% active solution of trimethyl hexadecyl ammonium chloride in a mixture of water and ethanol (levelling agent))	
"Quatramine C16/50"	4.5
PA57 silicone glycol copolymer levelling agent	0.5
Epoxy acrylate (Ex 20)	0.08
Component Compon	Raits by Weight

Viscosity Brookfield RVT #4/100/25°C = 460 cps.

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The composition was drawn down to a 5 micron coating on printed paper stock and cured using a mercury UV lamp of 80 w/cm at half power (ie 40 w/cm) half speed of 35 meters per min to provide a UV lamp energy of 25 millijoules per cm². The radiometer used to measure energy was a EIT Uvicure Radiometer model M365. The composition of the invention cured without addition of photoinitiator.

A conventional UV curable high gloss overprint varnish was prepared with the following composition.

19

PCT/AU00/00197

Comparative Example A

	7. 1
Craynor CN104 A80 = Epoxy Acrylate cut with 20% T.P.G.D.A.	43.0
T.M.P.E.O.T.A. = Ethoxylated Trimethylolpropane Triacrylate	48.5
Methyldiethanolamine = Amine synergist	3.0
Benzophenone, Photoinitiator	3.0
Irgacure 184 = 1-Hydroxy-cyclohexyl-phenyl-ketone, Photoinitiator	2.0
PA57 = Silicone glycol Copolymer (levelling and slip agent)	0.5
	100.0

Viscosity = Brookfield RVT #4/100/25°C = 460 cps

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The prior art composition required 5% photoinitiator and 3% amine synergist to cure under the same conditions.

Those skilled in the art will appreciate that there may be many variations and 10 modifications of the configuration described herein which are within the scope of the present invention.

ABSTRACT

Ultraviolet light curable resin compositions and methods of preparing radiation curable compositions are provided by this invention in which an amine adduct is formed by reaction of a secondary amine and epoxy and reacting the amine adduct with an unsaturated carboxylic acid to form a water soluble salt.

PCT/AU00/00197

Claims:

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A radiation curable resin composition comprising a water soluble amine
 salt prepolymer formed between an oligomer having at least one amine group and an unsaturated carboxylic acid.

- A resin composition according to claim 1 comprising an aqueous solution containing a weight ratio of amine salt prepolymer to water in the range of from 1:4 to 20:1.
 - 3. A resin composition according to claim 2 wherein said weight ratio is in the range of from 3:2 to 9:1.
- 4. A resin composition according to claim 1 which is curable by UV and contains less than 0.5% by weight based on the weight of the resin component of a UV initiator.
- 5. A resin composition according to claim 1 which is curable by UV in the absence of photoinitiator.
 - 6. A resin composition according to claim 1 wherein the oligomer having at least one amine group is selected from the group consisting of epoxy amine adducts, urea formaldehyde resins, melamine formaldehyde resins, amine-polyisocyanate adducts and michael adducts of an aliphatic amine and polyacrylate and/or polymethacrylate compound.
 - 7. A resin composition according to claim 1 wherein the oligomer having at least one amine group is an epoxy-amine adduct formed between a secondary amine and an epoxide of formula I:

$$CH_2$$
 CH_2 CH_2

PCT/AU00/00197

wherein n is from 0 to 10 and A is a diredical selected from the group consisting of aliphatic, substituted aliphatic, aromatic, substituted aromatic, cycloaliphatic, substituted cycloaliphatic, alkylene dicarbonyl and the diradical of formula II:

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$$R$$
 II

wherein R is an alkylene group of 1 to 4 carbon atoms.

- 10 8. A resin composition according to claim 7 or claim 8, wherein the amine adduct is formed using a molar ratio of amine to epoxide compound in the range of from 0.5:1 to 2.05:1.
- 9. A resin composition according to claim 7 or claim 8 wherein the molar ratio of amine to epoxide compound is 0.95:1 to 1.6:1.
 - 10. A resin composition according to claim 1 wherein the oligomer having at least one amine group is prepared from melamine, formaldehyde and/or glyoxal and optionally one or more alcohols selected from the group consisting of C_1 to C_6 alkanols, C_1 to C_6 alkyl ethers of C_1 to C_6 alkylene glycols and C_1 to C_6 alkylene glycols.
 - 11. A resin composition according to claim 1 wherein the oligomer having at least one amine group is a urethane-amine adduct prepared by reaction of a secondary amine with a polyisocyanate.
 - 12. A resin composition according to claim 1 wherein the urethane amine adduct is formed from a molar ratio of amine to polyisocyanate of from 0.90:1 to 1.6:1.

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13. A resin composition according to claim 1 wherein the oligomer having at least one amine group is a Michael adduct formed between a secondary amine and an unsaturated compound selected from the group consisting of monomers

and prepolymers comprising a plurality of unsaturated groups including at least one acrylate or methacrylate group.

14. A resin composition according to any one of claims 1 to 9 wherein the5 water soluble salt is formed using an amine of formula IV:

$$R \longrightarrow N \longrightarrow R^1$$
 IV

wherein R and R¹ are independently selected from straight and branched chain aliphatic of up to 6 carbon atoms optionally substituted by hydroxy and mixtures thereof.

15. A resin composition according to claim 1 wherein the unsaturated acid is selected from the group consisting of acrylic acid, methacrylic acid, crotonic acid, itaconic acid, maleic acid, mesaconic acid, citraconic acid, sorbic acid, fumaric acid and mixtures of two or more thereof.

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- 16. A method of preparing a radiation curable composition comprising forming an amine adduct by reaction of a secondary amine with an epoxy and reacting the amine adduct with an unsaturated carboxylic acid to form a water soluble salt.
- 17. A method according to claim 15 wherein the amine adduct and unsaturated carboxylic acid are reacted in the presence of water to provide an aqueous solution of a salt formed between the epoxy amine adduct and unsaturated carboxylic acid.
- 18. A method according to claim 15 wherein the reaction of the secondary amine and epoxy is exothermic and the reaction mixture is diluted with water during the exotherm.

WO 00/55228 PCT/AU00/00197

- 19. A method according to any one of claims 15 to 17 wherein water is added to provide a concentration of from 20 to 95%.
- 20. Use of a composition according to any one of claims 1 to 14 as a surface coating or ink.
 - 21. The use according to claim 18 wherein the composition is applied at a thickness of up to 200 microns.